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Mechanism of host substrate acetylation by a YopJ family effector

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Abstract

Yersinia outer protein J (YopJ) family of bacterial effectors depends on a novel acetyltransferase domain to acetylate signaling proteins from plant and animal hosts. However, the underlying mechanism is unclear. Here, we report the crystal structures of PopP2, a YopJ effector produced by the plant pathogen *Ralstonia solanacearum*, in complex with inositol hexaphosphate (IP₆), acetyl-coenzyme A (AcCoA), and/or substrate RRS1-R_{WRKY}. PopP2 recognizes the WRKYGQK motif of RRS1-R_{WRKY} to position a targeted lysine in the active site for acetylation. Importantly, the PopP2 – RRS1-R_{WRKY} association is allosterically regulated by IP₆ binding, suggestive of a previously unidentified role of the eukaryote-specific cofactor in substrate interaction. Furthermore, we provide evidence for the reaction intermediate of PopP2-mediated acetylation, an acetyl-cysteine covalent adduct, lending direct support to the “Ping-Pong”-like catalytic mechanism proposed for YopJ effectors. Together, our study provides critical mechanistic insights into the virulence activity of YopJ class of acetyltransferases.

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Author Contributions

Z.-M. Z. and L. G. determined the crystal structures of PopP2 complexes and conducted ITC assays. J.S. and L. G. performed NMR analysis. K.-W. M. and S.S. performed *in vitro* acetylation assays and *in vivo* functional analyses. Z.H. performed computational analysis. W.M. and J.S. designed and organized the study, Z.-M. Z., K.-W. M., W.M. and J.S. prepared the manuscript.

Competing Financial Interests

The authors declare no competing financial interest.

Introduction

The co-evolutionary arms race between pathogens and hosts has cultivated multiple lines of invasion and counter-invasion schemes. The first line of host defense involves the recognition of microbe-associated molecular patterns (MAMPs), which effectively restricts the colonization of the majority of potential pathogens^{1,2}. To counteract this MAMP-triggered immunity (MTI), successful pathogens employ secretion systems to inject virulence proteins, collectively known as effectors, into the host cells^{3,4}. Effectors secreted by the bacterial type III secretion system (T3SEs) play a central role in the pathogenesis of Gram-negative bacteria^{5,6}. Once translocated into the host cells, T3SEs manipulate specific host cellular processes, thereby promoting bacterial invasion and proliferation⁷. On the other hand, plant hosts have developed the nucleotide-binding/leucine-rich-repeat receptors (NLRs) to sense the activities of T3SEs, which consequently initiates a robust innate immune response termed effector-triggered immunity (ETI)³.

The YopJ superfamily of T3SEs define a broad class of novel acetyltransferases, with members produced from both animal and plant pathogens^{8,9}. YopJ effectors produced by mammalian pathogens, including YopJ from *Yersinia spp.*, AvrA from *Salmonella enterica* and VopA from *Vibrio parahaemolyticus*, acetylate specific serine, threonine and/or lysine residues in mitogen-activated protein kinase kinases (MAPKKs) to inhibit enzymatic activation^{10–13}, or ATP binding¹⁴, leading to suppressed inflammatory response^{10–14}. YopJ effectors produced by plant pathogens are more diversified on targeting host proteins⁸. Among them, PopP2 produced by the root-infecting bacterium *Ralstonia solanacearum* acetylates a conserved lysine residue in the heptad sequence WRKYGQK found in the defense-related WRKY transcription factors to inhibit their DNA binding activity^{15,16}. As such, PopP2 dampens the activation of defense genes in plant hosts. As a counter strategy, resistant plants evolved a chimeric NLR known as RRS1-R (Resistance to *Ralstonia solanacearum* 1), which incorporates a C-terminal DNA-binding WRKY domain as a “trap”. Acetylation of the WRKY domain of RRS1-R (RRS1-R_{WRKY}) by PopP2 activates ETI and confers resistance to *R. solanacearum*^{15,16}. By contrast, another YopJ homologue HopZ1a from *Pseudomonas syringae* modifies multiple substrates including tubulin¹⁷ and the Jasmonate ZIM domain (JAZ) protein¹⁸.

YopJ effectors bear no resemblance with other acetyltransferases. Instead, they depend on a catalytic core that is homologous to ubiquitin-like proteases (ULPs), with a conserved His/Cys/Glu or His/Cys/Asp catalytic triad^{19,20}. Furthermore, the activities of YopJ family effectors are regulated by a host cofactor inositol-hexakisphosphate (IP₆)^{20–23}, which is highly abundant in eukaryotes but absent in bacteria. Our recent study revealed that HopZ1a evolved a regulatory domain that, together with the ULP-like catalytic core, forms the IP₆- and AcCoA-binding pockets. IP₆ binding induces a conformational transition of HopZ1a to permit the formation of an AcCoA-binding pocket, thereby enhancing the acetylation reaction allosterically²⁰. However, the mechanism by which YopJ effectors bind and acetylate substrates remains unclear.

In this study, we determined the crystal structures of PopP2 in complex with IP₆, AcCoA and/or RRS1-R_{WRKY}. The complex structures define the interaction interface required for

the acetylation of RRS1-R and possibly WRKY transcription factors by PopP2. Importantly, the interaction of PopP2 with RRS1-R_{WRKY} is allosterically regulated by IP₆ and AcCoA, establishing a link between its cofactor binding and substrate recognition. Moreover, we characterized the structural details of the acetyl-cysteine reaction intermediate, which lends the long-sought-after evidence for the previously proposed “Ping-Pong” route of catalysis²⁴. Together, these studies provide key mechanistic insights into the virulence activity of PopP2 and other YopJ family effectors.

Result

Overall structures of PopP2 in complex with IP₆, AcCoA and/or RRS1-R_{WRKY}

To understand how PopP2 binds and acetylates host substrates, we solved the crystal structures of the acetyltransferase domain of PopP2 (PopP2_{WT}, residues 149–488) or its catalytic mutant C321A (PopP2_{C321A}) in complex with IP₆, AcCoA and/or the RRS1-R_{WRKY} domain (residues 1195–1273 of RRS1-R) (Fig. 1a–d, Supplementary Table 1). The structures of the PopP2_{WT} – IP₆, PopP2_{C321A} – IP₆ – AcCoA and PopP2_{WT} – IP₆ – AcCoA – RRS1-R_{WRKY} complexes reveal that PopP2 is comprised of a catalytic core assuming a ULP-like fold packed against a regulatory domain formed by the N- and C-terminal flanking sequences (Fig. 1b–d). The closely packed catalytic core and regulatory domain together form one IP₆-binding pocket (Fig. 1b), one AcCoA-binding cleft (Fig. 1c), and one surface groove cradling the RRS1-R_{WRKY} domain (Fig. 1d), similar to what was observed for HopZ1a (Fig. 1e). Structural comparison of all three PopP2 complexes reveals no substantial conformational change in PopP2, except for a disorder-to-order transition of loop_{β8β9} upon its binding to RRS1-R_{WRKY} (discussed in detail below) (Fig. 1b–d).

Despite with only ~30% sequence identity (Supplementary Fig. 1), PopP2 and HopZ1a superimpose well in their catalytic core and the IP₆- and AcCoA-binding pockets (Fig. 1e), giving a root-mean-square-deviation of 2.0 Å over the backbone of 224 residues. Their major structural difference lies in the linker sequence preceding the last β strand of the ULP fold, which forms a strand (β6) in PopP2 but assumes two helices in HopZ1a, as well as the N- and C-terminal helices that constitute one half of the regulatory domain (Fig. 1e). Another notable structural difference arises from their active sites: The active site of PopP2 is surrounded by several bulky hydrophobic residues (L191, L281 and F318), and T229 and S259, forming a deep cavity harboring the target lysine K1221; by contrast, the active site of HopZ1a is surrounded by two serines (S87 and S213), resulting in a shallow concave (Supplementary Fig. 2). Such distinct active site environments may explain why PopP2 prefers lysine for acetylation whereas HopZ1a acetylates serine and threonine residues²¹.

The RRS1-R_{WRKY} domain is comprised of a five-stranded antiparallel β-sheet stabilized by a zinc cluster, reminiscent of what was observed for other WRKY domains²⁵. The positively charged β2'- and β3'-strands of RRS1-R_{WRKY} penetrate into an acidic surface groove of PopP2, with the side chain of K1221 inserting into the active site (Fig. 1d and Supplementary Fig. 3). Note that each asymmetric unit of the crystal contains two PopP2_{WT}–IP₆–AcCoA–RRS1-R_{WRKY} complexes, crystal packing of which resulted in a displaced β1'-strand from one of the RRS1-R_{WRKY} molecules and reduced electron density

for the loop connecting $\beta 1'$ and $\beta 2'$ (Supplementary Fig. 4). For the purpose of clarity, our structural analysis only focuses on the intact region of the complex.

IP₆ regulation of the AcCoA binding of PopP2

The IP₆- and AcCoA-binding pockets of PopP2 are located distantly at the interface of the regulatory domain and the catalytic core (Fig. 1c). The PopP2 – IP₆ association is mediated by a cluster of polar residues (R208, S327, K331, K383, H384, N407, K408, R416, R419, K453 and R460) and the six phosphate groups of IP₆ through hydrogen bonding and/or electrostatic interactions (Fig. 2a). Note that the previously identified auto-acetylation site of PopP2, K383²⁶, is unmodified in the crystal structure. On the opposite side, the AcCoA molecule is embedded in a surface cleft extending from the regulatory domain toward the catalytic triad of PopP2, with the pantetheine arm of AcCoA untraceable, likely due to its high flexibility (Fig. 1c,d). The adenine moiety of AcCoA is buried in a hydrophobic pocket surrounded by the residues of both the catalytic core (K316, and S317) and the regulatory domain (Q386, V426, I444 and F446); while the 2'-hydroxyl, 3'-phosphate and 5'-pyrophosphate groups are hydrogen bonded to T445, S389 and K442, respectively (Fig. 2b). In the presence of RRS1-R_{WRKY}, we observed additional hydrogen bonds formed between the 5'-pyrophosphate and PopP2 S432 on the loop_{88β9} (Supplementary Fig. 5), suggesting a stabilization of the PopP2 – RRS1-R_{WRKY} interaction by AcCoA.

We then performed Nuclear Magnetic Resonance (NMR) spectroscopy and Isothermal Titration Calorimetry (ITC) on the catalytic mutant PopP2_{C321A} to evaluate the effect of IP₆ binding on the PopP2 – CoA interaction. First, it is apparent that the addition of IP₆ led to appearance of additional NMR signals in the aliphatic proton region of PopP2_{C321}, suggestive of IP₆-induced structural ordering (Supplementary Fig. 6a). Consistently, ITC analysis revealed that the presence of IP₆ enhanced the binding of PopP2_{C321A} to CoA from an undetectable level to a dissociation constant (K_d) of 57.2 μ M. However, such an interaction was severely impaired when mutations of the IP₆-binding sites (K383R or R416E) or AcCoA-binding sites (K316A/S389A/F446A) were introduced (Fig. 2c and Supplementary Table 2).

Furthermore, *In vitro* acetylation assay revealed that the presence of IP₆ significantly increased the PopP2-mediated acetylation of RRS1-R_{WRKY}, whereas the IP₆- and AcCoA-binding mutations nearly or completely abolished the acetyltransferase activity (Fig. 2d and Supplementary Fig. 6b,c). To test the activity of the PopP2 proteins *in planta*, PopP2 was fused to the N-terminal 79 residues of another T3SEs effector AvrRpt2, which serves as an artificial effector delivery system by bacterial pathogen *Pseudomonas syringae* pv. tomato (*Pto*)²⁷. As expected, *Pto* strain D28E expressing wild-type PopP2 triggered programmed cell death in resistant *Arabidopsis thaliana* plants (eco. Nd-1), which produces RRS1-R (Fig. 2e). By contrast, *Pto*D28E expressing the mutants of PopP2 all failed to trigger cell death (Fig. 2e), confirming that these mutations resulted in the loss of biological function of PopP2. Note that these mutants showed similar protein stability (Supplementary Fig. 6d). Together, these studies suggest that IP₆ enhances the activity of PopP2 by increasing its binding with AcCoA, a mechanism similar to that observed for HopZ1a.

Structural basis for the PopP2 – RRS1-R_{WRKY} interaction

The association of RRS1-R_{WRKY} with PopP2 is mediated by the α D-helix and three loops (loop β _{1 α C}, loop β _{2 β 3} and loop β _{8 β 9}) of PopP2, and the β 2'- β 3' segment of RRS1-R_{WRKY} (Fig 1d and Fig 3a). Notably, the strictly conserved WRKYGQK motif of RRS1-R_{WRKY} (Supplementary Fig. 7a) is recognized by PopP2 through a number of hydrogen bonds and van der Waals contacts (Fig. 3b). Along the β 2'-strand, the side chain of RRS1-R_{WRKY} R1216 donates a water-mediated hydrogen bond to PopP2 E364; the backbone and side chain of RRS1-R_{WRKY} K1217 form hydrogen bonds with PopP2 D292 and N296, and E284, respectively; the aromatic ring of RRS1-R_{WRKY} Y1218 inserts into the cavity formed by PopP2 M219, M221 and D257, with its backbone and side chain engaging water-mediated hydrogen bonds with PopP2 D257, R262 and N296, and M221, D257 and E300, respectively (Fig. 3b); and the backbone of RRS1-R_{WRKY} Q1220 forms a water-mediated hydrogen bond with PopP2 Y293. Most notably, the side chain of RRS1-R_{WRKY} K1221, the target acetylation site, inserts into the catalytic pocket formed by PopP2 L191, S259, L281, Q315 and F318, with its side chain interacting with PopP2 H260, F318 and C321 through direct or water-mediated hydrogen bonds (Fig. 3a). Mutation of RRS1-R_{WRKY} K1221 into glutamine, mimic of acetylated lysine, led to abolished DNA binding activity of RRS1-R_{WRKY} (Supplementary Fig. 7b), supporting previous findings that PopP2-mediated acetylation disrupts the DNA binding activities of WRKY transcription factors^{15,16}. C-terminal to the WRKYGQK motif, the loop β 2' β 3' stacks against two loops (loop β 5 α E and loop β 8 β 9) of PopP2, with hydrogen bond interactions involving RRS1-R_{WRKY} D1222, I1223, G1225, S1226 and R1227 and PopP2 E184, R427, R429 and A430 (Fig. 3c). Additional intermolecular contacts are mediated by the β 3'-strand of RRS1-R_{WRKY} (Y1232, R1234, A1236, K1238 and F1239), which makes hydrogen bonding or van der Waals contacts with loop β _{1 α C} (V227 and T229), α D (D295, N296 and M299) and loop β 6 β 7 (E364 and L365) of PopP2 (Fig. 3b).

Interestingly, the majority of the PopP2-interacting residues is strictly conserved within the WRKY domains, and also required for DNA binding (Supplementary Fig 7a)²⁸. We therefore tested whether PopP2 binding would directly affect the DNA binding of RRS1-R_{WRKY} using electrophoretic mobility shift assay (EMSA). We observed that increasing amount of PopP2 led to a modest reduction in the DNA binding of RRS1-R_{WRKY} (Supplementary Fig. 7c), suggesting that the PopP2 interaction contributes to the decreased DNA binding activity of WRKY domains.

To evaluate how the interaction with the WRKY domain affects the virulence activity of PopP2, we generated an E284A/D292A/N296A triple mutant, which eliminates the PopP2–RRS1-R_{WRKY} binding (Supplementary Fig. 8 and Supplementary Table 2). Although this mutant retains the activities of CoA-binding (Supplementary Table 2) and auto-acetylation (Supplementary Fig. 9a), it was no longer able to acetylate RRS1-R_{WRKY} *in vitro*, regardless of the presence of IP₆ (Fig. 3d), and failed to induce cell death in the resistant *Arabidopsis* plants (Fig. 3e). These data suggest that the RRS1-R_{WRKY}-interacting region of PopP2 is essential for its biological function *in planta*.

Distinct substrate binding strategies between YopJ effectors and ULPs

To assess how YopJ family of effectors evolves into a class of acetyltransferases with diverse substrate specificities, we compared the substrate binding sites of PopP2 with the corresponding regions of HopZ1a. Notably, the WRKY-binding sites of PopP2 (loop $\beta_{1\alpha C}$, αD and loop $\beta_{8\beta 9}$) are largely preserved in HopZ1a, albeit with distinct conformations (Fig 4a–c), implying that this region may also underlie the substrate specificity of HopZ1a. Consistently, our previous study revealed that introduction of a mutation (C141K) to the HopZ1a region, equivalent to loop $\beta_{1\alpha C}$ of PopP2 (Fig 4b), leads to altered allelic specificity of HopZ1b, an allele evolved from a HopZ1a-like ancestor²⁹, suggesting that alteration in substrate binding might be an important mechanism for the evolution of YopJ effectors.

Next, we performed structural comparison between PopP2_{WT} – IP₆ – AcCoA – RRS1-R_{WRKY} and one of the closely related enzyme-substrate complexes, the ULP1-SMT3 complex³⁰ (Fig. 4d–f). ULPs contain four conserved regions for recognition of SUMO- or ubiquitin-like substrates (R1 and R2) or guiding the C-terminus of the substrate toward the catalytic site (R3 and R4)³¹ (Fig 4d–f and Supplementary Fig. 1). By comparison, although PopP2 and ULP1 contain similar catalytic sites (R3 and R4), the substrate-recognition regions in ULP1 are either absent (in the case of R1) or structurally divergent (in the case of R2) in PopP2 (Fig. 4d). Notably, the R2-corresponding region of PopP2 not only blocks R3, but also contributes to the formation of a surface groove right on top of the catalytic center, resulting in the distinct substrate-recognition site for PopP2 (Fig. 1d and Fig. 4e). In fact, the R1- and R2-corresponding regions are also structurally varied in HopZ1a (Fig. 1e), further highlighting the functional divergence between YopJ effectors and ULPs.

IP₆ regulation of the PopP2 – WRKY interaction

The observation that the loop $\beta_{8\beta 9}$ of PopP2 is involved in binding to both AcCoA and RRS1-R_{WRKY} (Fig. 1d) suggests a role of this region in linking cofactor binding to substrate binding. Indeed, structural comparison of the PopP2_{WT} – IP₆, PopP2_{C321A} – IP₆ – AcCoA and PopP2_{WT} – IP₆ – AcCoA – RRS1-R_{WRKY} complexes reveals that the β_8 - β_9 segment of PopP2 is partly stabilized by AcCoA binding, and becomes even more ordered upon RRS1-R_{WRKY} binding (Fig. 5a,b). These observations, together with the observation that the loop $\beta_{8\beta 9}$ -equivalent region in HopZ1a undergoes an IP₆-induced structural ordering²⁰, prompts us to investigate whether the IP₆ or AcCoA binding regulates the PopP2 – RRS1-R_{WRKY} association. Indeed, whereas the binding affinity of IP₆-free PopP2 for RRS1-R_{WRKY} was undetectable, addition of IP₆ increased the binding to a K_d of 11.0 μ M, and another ~3.3-fold increase when CoA was also added (Fig. 5c and Supplementary Table 2). By contrast, the IP₆-binding deficient mutations (K383R and R416A) largely abolished the interaction (Supplementary Fig. 8 and Supplementary Table 2). Consistently, molecular dynamics (MD) simulations revealed that the root mean square fluctuation (RMSF) of the atomic fluctuations of PopP2 loop $\beta_{8\beta 9}$ undergoes a gradual reduction upon binding to IP₆, IP₆ and AcCoA, or IP₆, AcCoA and RRS1-R_{WRKY}, suggesting that IP₆ and AcCoA regulate the interaction of PopP2 with RRS1-R_{WRKY} by inducing structural ordering of loop $\beta_{8\beta 9}$ (Fig. 5d). Taken together, these data establish that the IP₆ binding not only allosterically stimulates the formation of AcCoA-binding pocket, but also regulates the substrate binding of PopP2.

Structure of a PopP2 – AcCoA covalent adduct

Given the fact that YopJ effectors contain a ULP-like catalytic triad, it has been proposed that they acetylate substrates via a “Ping-Pong” route of catalysis¹², but not the “direct transfer” mechanism adopted by the GCN5-related N-acetyltransferase (GNAT) or the MYST family of acetyltransferases^{32–34}. The “Ping-Pong” model predicts the presence of an acetyl-cysteine intermediate on the reaction pathway, generated by nucleophilic attack of the catalytic cysteine on AcCoA, which then reacts with the target residue to create an acetylated product^{12,35}. To provide experimental support for this model, we captured the reaction intermediate between PopP2 and AcCoA by soaking AcCoA into the crystals of the IP₆-bound, enzymatically active mutant PopP2_{F318S} (Supplementary Fig. 9b). In comparison with wild-type PopP2, this mutant yielded a higher-quality crystal structure.

The overall structure of the PopP2_{F318S} – IP₆ – AcCoA complex is identical to that of the PopP2_{C321A} – IP₆ – AcCoA complex (Fig. 6a), except for the presence of an acetyl group covalently attached to the side chain of C321 (SCY321), surrounded by residues S259, H260, Q315, F319 and R322 (Fig. 6a). A water molecule, situated in a position that is otherwise occupied by the side chain amino group of the target lysine, bridges the acetyl group with H260 of the catalytic triad (Fig. 6a, b). Meanwhile, the water molecule also receives a hydrogen bond from another highly conserved residue, Q315 (Fig. 6a and Supplementary Fig. 1). The presence of a target lysine would presumably lead to eviction of this water molecule (Fig. 6b). Subsequently, hydrogen bonding with both H260 and the acetyl-cysteine adduct would permit the target lysine to initiate a nucleophilic attack of the acetyl-cysteine, resulting in substrate acetylation. Consistently, mutation of H260 into alanine leads to abolished enzymatic activity of PopP2 as well as PopP2-triggered cell death in *Arabidopsis* (Supplementary Fig. 9b,c), reinforcing the notion that the H260-equivalent residue serves as the catalytic base in YopJ effector-mediated acetylation.

Discussion

Enzyme-substrate interaction of PopP2 determines the outcome of pathogen invasion

The molecular interactions between bacterial effectors and host targets critically influence the outcome of pathogenic infections. Although PopP2 has evolved to target multiple WRKY transcription factors to suppress MTI, the incorporation of a decoy WRKY domain into the C-terminus of RRS1-R in resistant *Arabidopsis* plants successfully translates the virulence activity of PopP2 into defense activation^{15,16}. This study provides the structural basis for the recognition of the conserved WRKYGQK motif by PopP2, which guides the insertion of the target lysine into the active site pocket. Strikingly, the IP₆-induced structural ordering of loop_{88β9} enhances both AcCoA and WRKY bindings, providing a novel mechanism by which IP₆ regulates the basal activity of PopP2. Given the structural similarity between PopP2 and HopZ1a, it is conceivable that such an IP₆ regulation may help ensure efficient host-specific substrate recognition of both PopP2 and HopZ1a in plant cells.

Structural evolution and target recognition of YopJ effectors

YopJ effectors modify a diverse array of host proteins to promote pathogenesis⁸. Structural analysis of the PopP2_{WT} – IP₆ – AcCoA – RRS1-R_{WRKY} complex suggests that YopJ effectors have evolved a distinct substrate-recognition strategy from that of ULPs. Furthermore, it provides insight into how YopJ effectors achieve distinct substrate specificities. In particular, the WRKY-binding sites of PopP2 (e.g. αD-helix), despite with structural homology with the corresponding region of HopZ1a, manifest substantial conformational and sequence divergence from other YopJ effectors (Supplementary Fig. 1), implying that the substrate binding sites of YopJ effectors are under strong selective pressure during arms race with specific hosts. Taken together, this study provides a framework for further understanding of the effector-target interaction of other YopJ effectors.

YopJ family effectors share a novel acetyltransferase mechanism

The finding that YopJ effectors contain a conserved ULP-like catalytic core challenges the current structure-function paradigm of acetyltransferases. Here, structural characterization of the acetyl-cysteine reaction intermediate of PopP2 lends a key support to a previous notion that YopJ effector-mediated acetylation mechanistically resembles the proteolysis process mediated by cysteine proteases²⁴; that is, YopJ effectors employ a “Ping-Pong”-like route of catalysis involving sequential binding of AcCoA and protein substrates. It is also worth pointing out that the presence of the PopP2 – IP₆ – AcCoA – RRS1-R_{WRKY} quaternary complex also suggests that the substrate binding of YopJ effector may precede a complete release of the product CoA (Fig. 6c), which deviates from the classic “Ping-Pong” model.

Materials and Methods

Protein Preparation

The DNA sequences encoding PopP2, except for its N-terminal 148 residues that are predicted to be structurally disordered and responsible for transferring the effector into host cells by *Ralstonia solanacearum*, and RRS1-R_{WRKY} (residues 1195–1273) were inserted in frame with a hexahistidine-SUMO tag in the pRSF-Duet vector. The recombinant proteins of PopP2 or RRS1-R_{WRKY} were overexpressed in BL21(DE3) RIL cell strain overnight by the addition of 0.4 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) at 20°C when the cell density reached an OD₆₀₀ of 0.8. For Se-Met labeling, cells cultured in M9 minimum medium to mid-log phase were induced for 16 hours after the addition of 50 mg/L of each of the following amino acids: lysine, phenylalanine, threonine, isoleucine, leucine, valine and L-selenomethionine. Cells were lysed in buffer containing 50 mM Tris-HCl (pH 8.0), 1 M NaCl, 25 mM imidazole, 0.5 mM DTT and 1 mM PMSF. The soluble fraction was subjected to a Nickel column. For purification of PopP2, the eluted fusion proteins were cleaved by ULP1, followed by further purification using hydrophobic interaction chromatography (Phenyl HP, GE Healthcare). Peak fractions were collected, concentrated and subsequently applied to Superdex 200 16/600 (GE Healthcare) pre-equilibrated with a buffer containing 25 mM Tris (pH 7.5), 200 mM NaCl, 5 mM IP₆, 2 mM DTT and 5% glycerol. Purified PopP2 samples were stored in –80°C at a concentration of ~50 mg/mL for future use. For purification of RRS1-R_{WRKY}, the RRS1-R_{WRKY} protein in fusion with the SUMO tag was cleaved by ULP1. Subsequently, the RRS1-R_{WRKY} protein was purified through a Heparin

column (GE Healthcare) followed by size exclusion chromatography on a Superdex 200 16/600 column pre-equilibrated with a buffer containing 25 mM Tris-HCl (pH 7.5), 100 mM NaCl and 2 mM DTT.

Crystallization and structure determination

The PopP2_{C321A} – IP₆ and PopP2_{WT} – IP₆ complexes, prepared by mixing of the corresponding proteins with IP₆ in a molar ratio of 1:10, was crystallized by hanging-drop diffusion method. Crystals appeared in a buffer containing 0.1 M HEPES (pH 7.5), 20% (v/v) PEG4000 and 10% (v/v) glycerol. The PopP2_{F318S} – IP₆ complex was crystallized in a buffer containing 0.1 mM imidazole (pH 8.0), 30% (v/v) PEG8000, 200 mM NaCl and 4% (v/v) isopropanol. The PopP2_{WT} – IP₆ – RRS1-R_{WRKY} complex was prepared by mixing wild-type PopP2 and RRS1-R_{WRKY} in a molar ratio of 1:1.1 in a buffer containing 25 mM Tris-HCl (pH 7.5), 250 mM NaCl, 50 mM Arg/Glu, 5% glycerol and 2 mM IP₆, and crystallized in a buffer containing 0.1 mM Tris-HCl (pH 8.5), and 20% ethanol. The crystals for the PopP2_{C321A} – IP₆ – AcCoA, PopP2_{F318S} – IP₆ – AcCoA and PopP2_{WT} – IP₆ – AcCoA – RRS1-R_{WRKY} complexes were obtained by soaking crystals in the corresponding crystallization buffer supplemented with 5 mM AcCoA for overnight before flash frozen in liquid nitrogen. Crystallization buffer was supplemented with glycerol or PEG400 to a final concentration of 15–20% (v/v) as cryo-protectant before flash freezing crystals in liquid nitrogen. Diffraction data were collected on beamline 5.0.1 and 5.0.2 at the Advanced Light Source (ALS), Lawrence Berkeley National Laboratory and 24-IDE NE-CAT beamline at the Advanced Photo Source (APS), Argonne National Laboratory. The data were indexed, integrated and scaled by HKL2000 package³⁶ or XDS³⁷. The structure of PopP2_{C321A}–IP₆ complex was solved by the Se-SAD method and the initial model was built by the Autosol module embedded in the PHENIX software package³⁸. Iterative cycles of model rebuilding and refinement were performed using COOT³⁹ and PHENIX, respectively. The structures of the PopP2_{WT} – IP₆, PopP2_{C321A} – IP₆ – AcCoA, PopP2_{F318S} – IP₆ – AcCoA and PopP2_{WT} – IP₆ – AcCoA – RRS1-R_{WRKY} complexes were solved by molecular replacement using PHASER⁴⁰ with PopP2_{C321A} – IP₆ and/or WRKY1 (PDB 2AYD) structure as searching model.

The statistics for data collection and structural refinement for the PopP2 complexes is summarized in Supplementary Table 1.

ITC measurements

IP₆ and AcCoA were dissolved in water at an initial concentration of 100 mM (adjusted to pH 7.5 by 0.1 mM NaOH) and 30 mM, respectively, and diluted to 1 mM by the ITC buffer before the titration. Protein samples for the RRS1-R_{WRKY} domain, wild-type, and mutant PopP2 were dialyzed against the ITC buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 0.2 mM IP₆ and 5% glycerol) at 4°C overnight. A MicroCal iTC200 system (GE Healthcare) was used to conduct the ITC measurements. A total of 17–20 injections with a spacing of 180 s and a reference power of 5 µcal/s were performed at 25°C. The ITC curves were processed with software ORIGIN (MicroCal) using one-site fitting model.

In vitro acetylation assay

In vitro acetylation assays were carried out to examine the acetyltransferase activity of PopP2. For [^{14}C] labeling experiments, 4 μg PopP2 were incubated with [^{14}C]-acetyl-CoA (55 $\mu\text{Ci}/\mu\text{mol}$) in 25 μL reactions (50 mM HEPES (pH 8.0), 150mM NaCl, 1 mM DTT, 1 mM PMSF, 10 mM sodium butyrate and 5% glycerol) at room temperature for an hour. 22.5 μg SUMO-RRS1-R_{WRKY} was used as the substrates of PopP2. 10 nM IP₆ and 30 μM AcCoA were added. The reactions were stopped by 4 \times Laemmli buffer and subjected to 12% SDS-PAGE. Acetylated proteins were detected after five days. After autoradiography, the protein gels were stained with Coomassie Blue as a loading control.

For immunoblot experiments, similar experiments were carried out except that 60 μM ^{12}C -acetyl-CoA was used instead. Acetylated proteins were detected using Ac-K antibody (Cell signaling, Ac-K-103). The protein gels were stained with Coomassie Blue as a loading control

Pseudomonas syringae infection assays

DNA sequence encoding N-terminal AvrRpt2 (residue 1–79) fused with truncated wild-type and mutant PopP2 carrying a C-terminal HA tag were cloned into the vector pUCP20tk. The genes are under the control of native *avrRpt2* promoter. The plasmids were transformed into *Pseudomonas syringae* pv. tomato strain D28E (*Pto*D28E), which is an effector-less mutant of wild-type strain DC3000. The expression and stability of the chimeric proteins in *Pto*D28E was confirmed by immunoblotting using an anti-HA antibody (Roche Diagnostics) after the cells were induced in MG minimal medium for two days. Secretion of the chimeric proteins in plant cells during bacterial infection was not checked.

For hypersensitive response assay, *P. syringae* strains were grown on mannitol-glutamate (MG) agar at room temperature for two days. Leaves of five-week-old *Arabidopsis thaliana* plants (eco. Nd-1) were infiltrated with bacterial suspensions at OD₆₀₀ = 0.4 (approximately 4×10^8 cfu/mL) in 10 mM MgSO₄. The inoculated plants were transferred to a growth chamber (22°C and 16/8 light/dark regime, 90% humidity). Cell death symptoms were monitored at 24 hours post inoculation (hpi).

Molecular dynamics (MD) simulations

Protein structure was prepared with AmberTools16⁴¹. The pantetheine arm of AcCoA was interactively docked using YetiX⁴² using the solved crystal structure as template. Each complex was explicitly solvated in a rectangular box of TIP3P model water (at least 12 Å from the complex to avoid periodic artifacts from occurring) by using AMBER16 with the ff14SB and gaff2 force field for protein and ligands. Ligand partial charges were calculated by RESP fit with Gaussian09 using the B3LYP/6–31G* basis set after geometry optimization. The Zinc metal center parameters of the RRS1-R_{WRKY} domain were processed with ZAFF parameters⁴³. For each system, 200ns production MD simulations were performed for trajectory analysis. The last 50ns trajectory were subjected to RMSF analysis of C-alpha coordinates by cpptraj⁴⁴.

Electrophoretic mobility shift assay

The samples were prepared by mixing various amounts of PopP2, RRS1-R_{WRKY} domain and a 16 mer W-box DNA duplex (upper strand: CGCCTTTGACCAGCGC) in a total volume of 40 μ L of solution containing 25 mM Tris-HCl (pH7.5), 200 mM NaCl and 5 mM DTT. For titration of PopP2, the sample mixture was supplemented with 2 mM IP₆ and 1.5 mM AcCoA, and incubated on ice for 1 hr before gel electrophoresis. The samples were loaded onto 10% non-denaturing polyacrylamide gel and electrophoresed at 4°C in 0.25 \times TBE buffer. The gel was stained by ethidium bromide and visualized under UV light.

Data availability

Coordinates and structure factors have been deposited in the Protein Data Bank under accession codes PDB 5W3T (PopP2_{WT} – IP₆), PDB 5W3Y (PopP2_{C321A} – IP₆ – AcCoA), PDB 5W40 (PopP2_{F318S}-IP₆-AcCoA) and PDB 5W3X (PopP2_{WT} – IP₆ – AcCoA – RRS1-R_{WRKY}).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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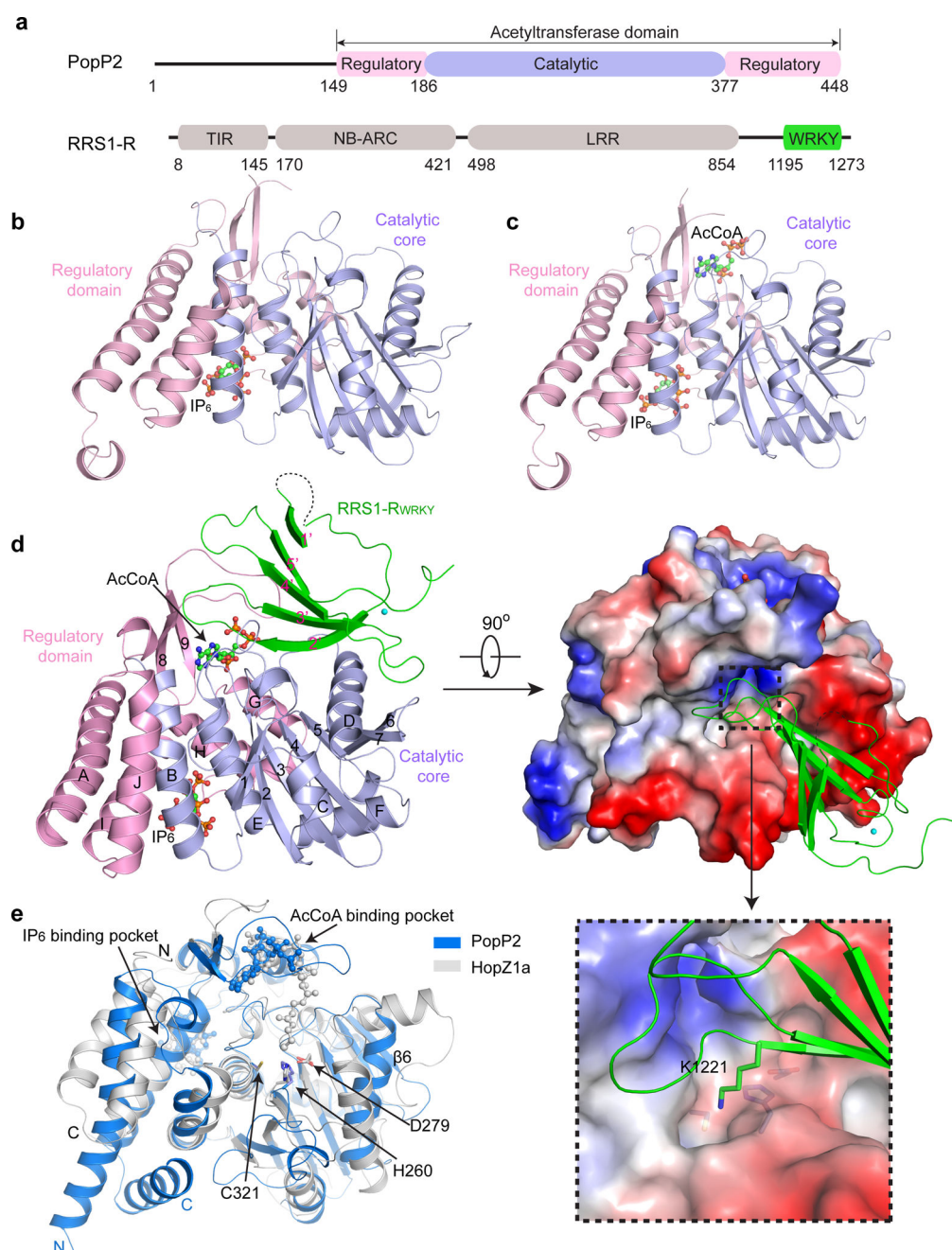


Figure 1. Crystal structures of PopP2 in complex with IP₆, AcCoA and/or RRS1-R_{WRKY}
a, Domain architectures of PopP2 and RRS1-R, with individual domains colored differently. TIR, Toll/interleukin1 receptor; NB, nucleotide binding; ARC, Apaf-1, R-protein and CED-4; LRR, leucine-rich repeats. Similar color schemes are used in other figures unless otherwise indicated. **b**, Crystal structure of PopP2_{WT} in complex with IP₆. **c**, Crystal structure of PopP2_{C321A} in complex with IP₆ and AcCoA. **d**, Ribbon and surface representations of the PopP2_{WT}-IP₆-AcCoA-RRS1-R_{WRKY} complex. The α-helices and β-strands in PopP2 are labeled from A to J and 1 to 9, respectively. The β-strands in RRS1-R_{WRKY} are labeled from 1' to 5'. The IP₆ and AcCoA molecules are shown in ball-and-stick

representation. The zinc ion is shown as a cyan sphere. The active site of PopP2 and the target lysine K1221 in RRS1-R_{WRKY} are shown in an expanded view. **e**, Structural alignment of PopP2 and HopZ1a, colored in blue and grey, respectively. The catalytic triad (H260/D279/C321 in PopP2 or H150/E170/C216 in HopZ1a) is shown in stick representation.

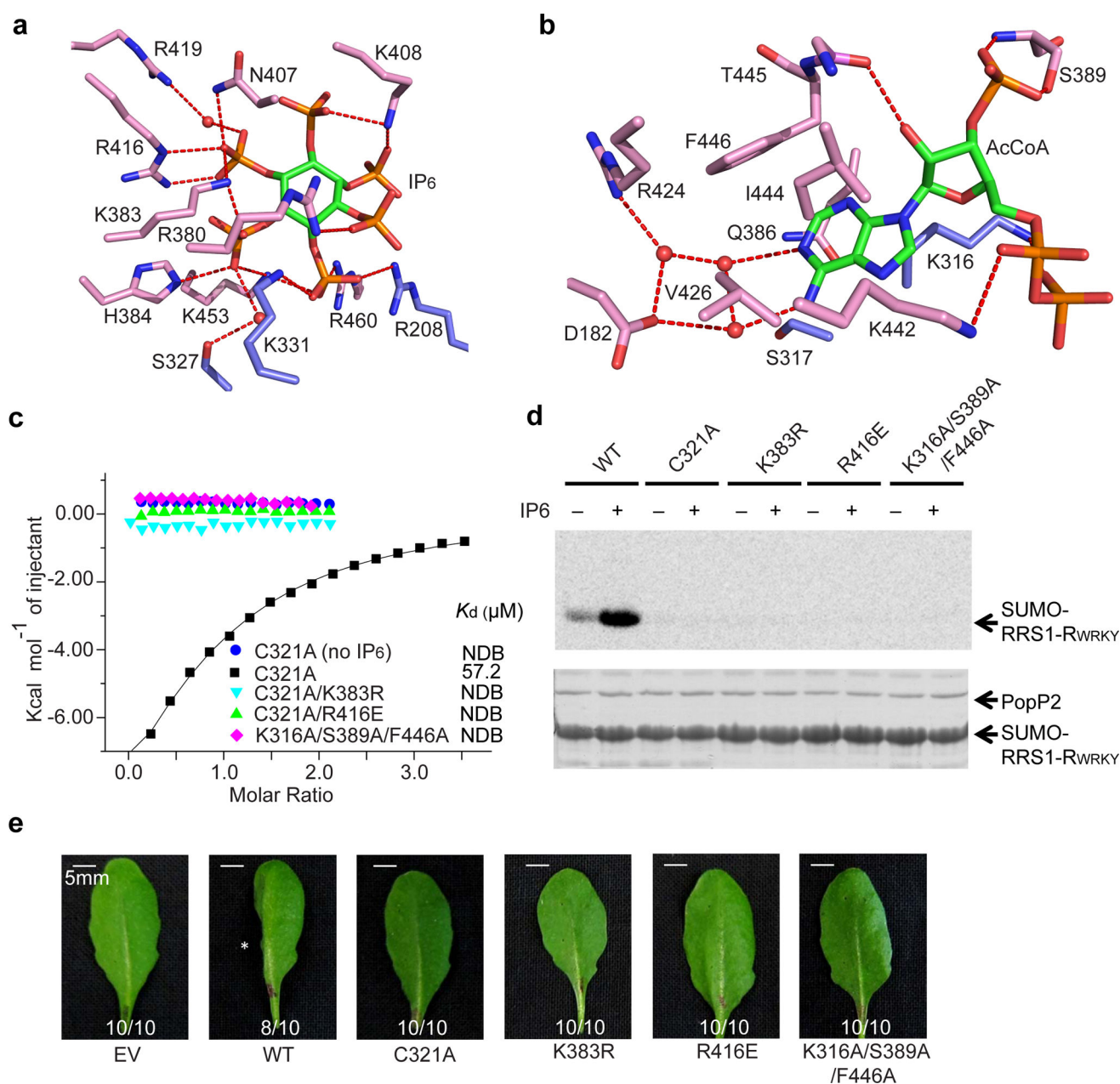


Figure 2. IP₆-dependent acetyltransferase activity of PopP2

a, Close-up view of the PopP2 – IP₆ interaction. **b**, Close-up view of the PopP2 – AcCoA interaction. In **a** and **b**, the ligands are shown as green sticks, the hydrogen bonds are depicted as red dashed lines, and the water molecules are shown as red spheres. **c**, ITC binding curves of PopP2 mutants with AcCoA in the presence or absence of IP₆. **d**, *In vitro* acetylation activity of wild-type (WT) or mutant PopP2. The acetylated proteins were detected by autoradiography (top), and the protein amount is indicated by Coomassie blue staining (bottom). **e**, Cell death-triggering activity of PopP2 in *Arabidopsis* eco. Nd-1. The asterisk indicates the cell death symptom triggered by wild-type PopP2. Ratios indicate number of leaves with indicated phenotype/total number of inoculated leaves in each

treatment. Experiments in **d** and **e** were repeated twice and once, respectively, with consistent results.

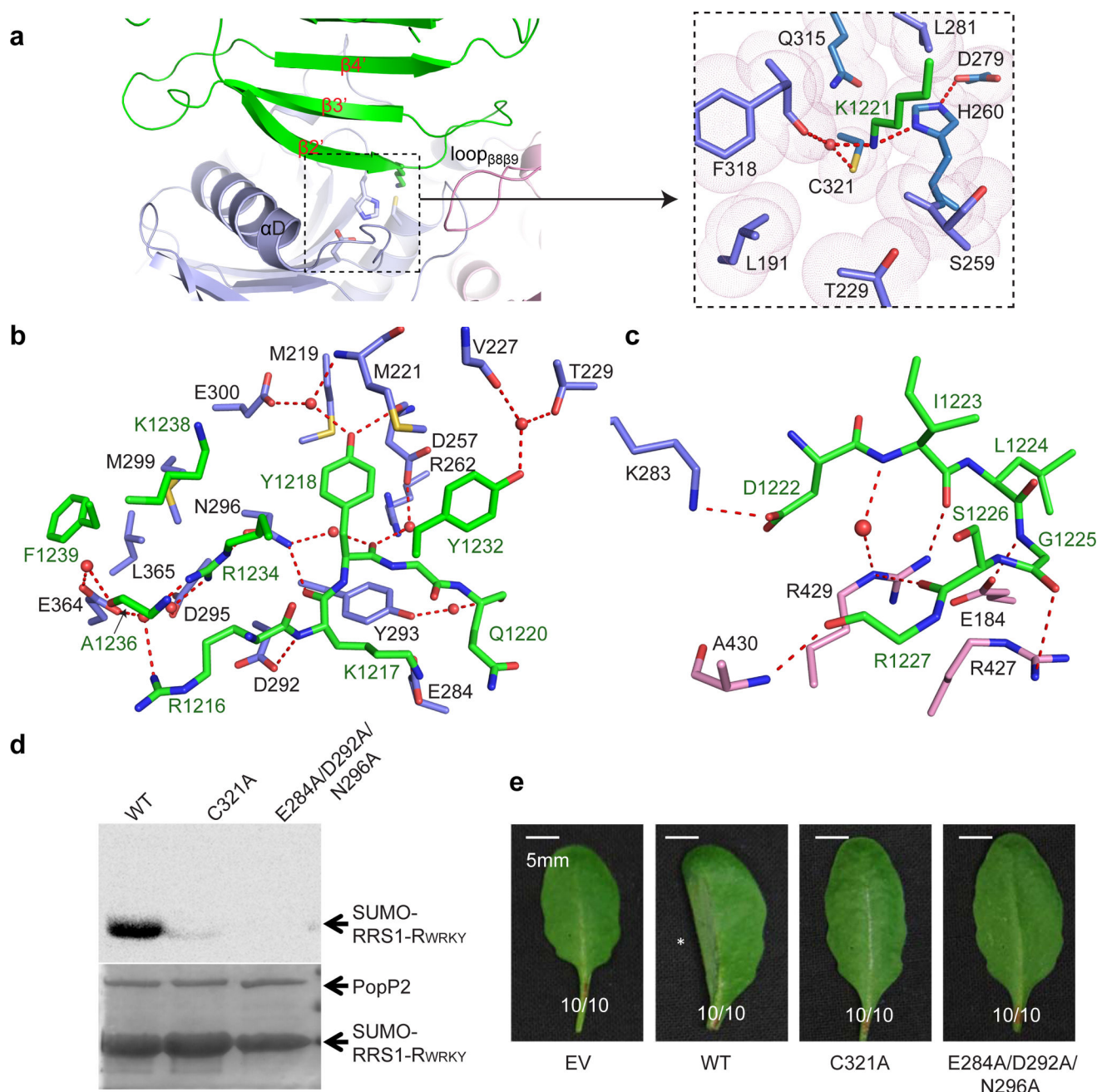


Figure 3. The PopP2 – RRS1-RWRKY interaction

a, The interface between PopP2 and RRS1-RWRKY, with the catalytic site shown in expanded view. **b**, Close-up views of the interactions of PopP2 with $\beta 2$, $\beta 3$ and Loop $\beta 3\beta 4$ of RRS1-RWRKY. **c**, Close-up views of the interactions of PopP2 with loop $\beta 2\beta 3$ of RRS1-RWRKY. Hydrogen bonds are depicted as dashed lines, and water molecules are shown as red spheres. **d**, *In vitro* acetylation activity of wild-type (WT) or mutant PopP2. The acetylated proteins were detected by autoradiography, and the protein amount is indicated by Coomassie blue staining. **e**, Cell death-triggering ability of PopP2 in *Arabidopsis* eco. Nd-1. The asterisk indicates the cell death symptom triggered by wild-type PopP2. Ratios indicate

number of leaves with indicated phenotype/total number of inoculated leaves in each treatment. Experiments in **d** and **e** were repeated twice and once, respectively, with consistent results.

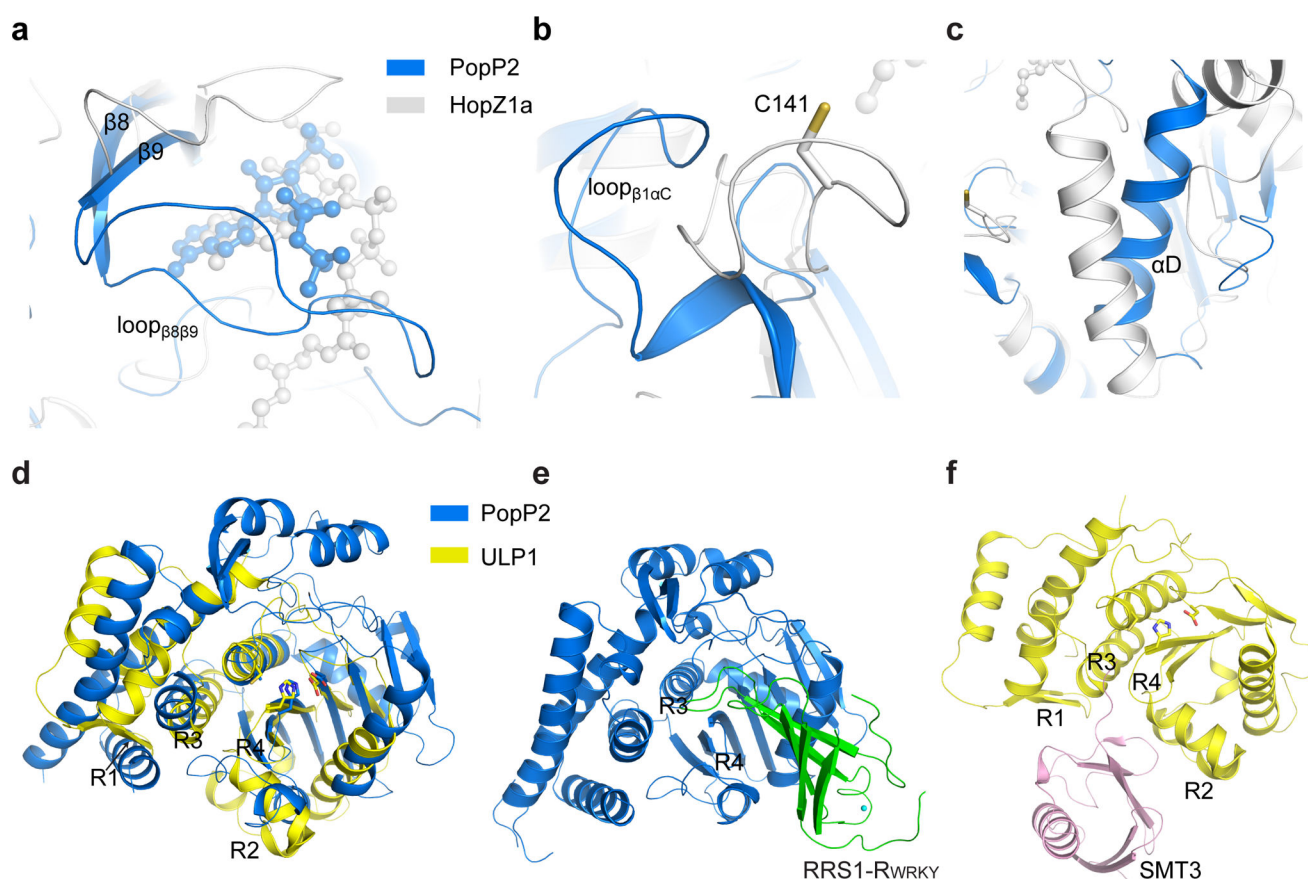


Figure 4. Structural comparison of PopP2_{WT} – RRS1-R_{WRKY}, HopZ1a and ULP1 – SMT3
a–c, Structural overlay of the substrate binding sites of PopP2 (blue) with the corresponding regions in HopZ1a (grey). **d**, Overlaid structures of PopP2 (blue) and ULP1 (PDB 1EUU, yellow), with the catalytic triads shown in stick representation. **e–f**, Crystal structures of the PopP2_{WT} – IP₆ – AcCoA – RRS1-R_{WRKY} **e** and ULP1 – SMT3 **f** complexes in the same orientation as shown in **d**. The substrate binding regions of ULP1, and the R3- and R4-equivalent regions in PopP2, are labeled.

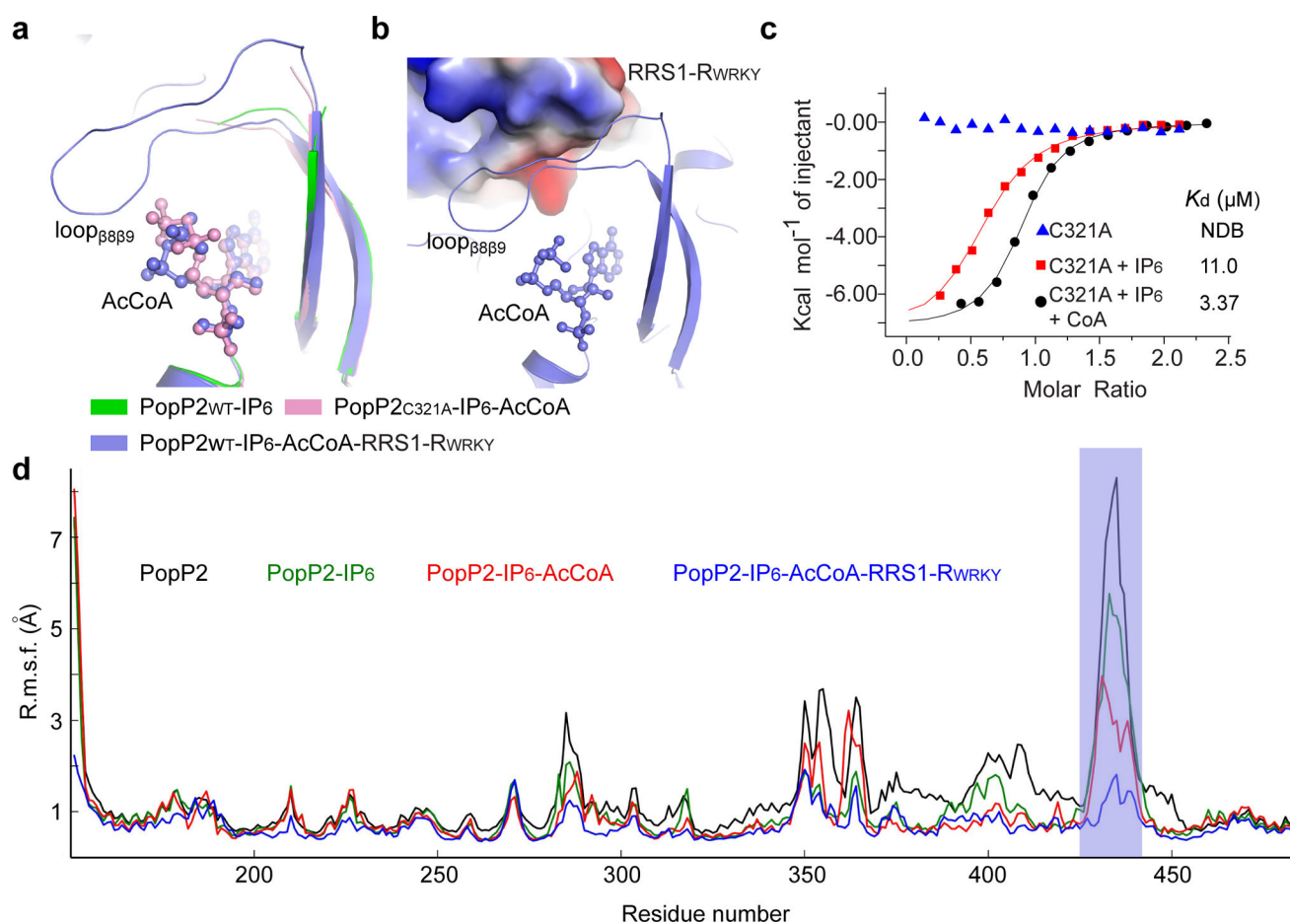


Figure 5. IP₆-stimulated binding between PopP2 with RRS1-R_{WRKY}

a, Close-up view of the conformational change of the $\beta 8$ - $\beta 9$ segment of PopP2 in different complexes. AcCoA is shown in ball-and-stick representation. **b**, The structure of loop $\beta 8\beta 9$ in the PopP2_{WT}-IP₆-AcCoA-RRS1-R_{WRKY} complex. The PopP2-binding site of RRS1-R_{WRKY} is shown in electrostatic surface representation. **c**, ITC binding curves of PopP2 with RRS1-R_{WRKY} in the presence or absence of IP₆ and/or CoA. **d**, RMSF of atomic fluctuations of PopP2, free or in complex with IP₆, AcCoA and/or RRS1-R_{WRKY}. The region corresponding to loop $\beta 8\beta 9$ is shaded in blue.

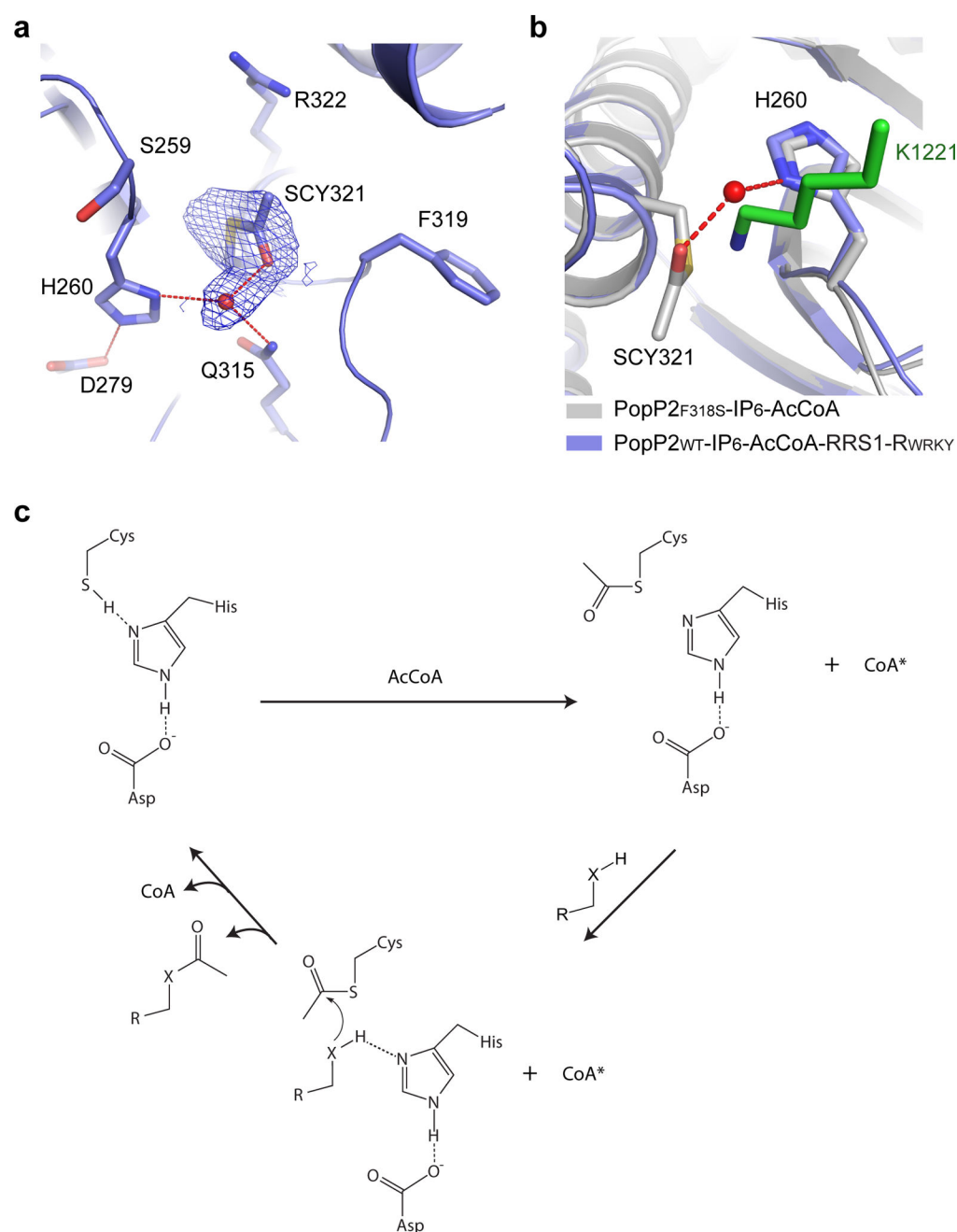


Figure 6. A model for enzymatic catalysis of YopJ effectors

a, Close-up view of the catalytic site in the crystal structure of PopP2^{F318S} – IP₆ – AcCoA, with the 2F_o–F_c omit map of the acetyl-cysteine (SCY321) contoured at 1.0σ level. The water molecule is shown as a red sphere and the hydrogen bonds are shown as dashed lines.

b, The active site of PopP2^{F318S} – IP₆ – AcCoA overlaid with that of PopP2^{WT}– IP₆ – AcCoA– RRS1–RWRKY. The target lysine is shown in green stick. **c**, A model for the catalytic steps of YopJ effector-mediated acetylation. *The adenosine moiety of CoA

remains bound to PopP2, whereas the pantetheine arm is released from the catalytic center. “X” denotes an NH₂ group or oxygen atom.

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